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NEWS 24 Feb 26 NTIS now allows simultaneous left and right truncation
NEWS 25 Feb 26 PCTFULL now contains images
NEWS 26 Mar 04 SDI PACKAGE for monthly delivery of multifile SDI results NEWS 27 Mar 19 APOLLIT offering free connect time in April 2003 NEWS 28 Mar 20 EVENTLINE will be removed from STN NEWS 29 Mar 24 PATDPAFULL now available on STN NEWS 30 Mar 24 Additional information for trade-named substances without structures available in REGISTRY NEWS 31 Apr 11 Display formats in DGENE enhanced NEWS 32 Apr 14 MEDLINE Reload NEWS 33 Apr 17 Polymer searching in REGISTRY enhanced NEWS 34 Apr 21 Indexing from 1947 to 1956 being added to records in CA/CAPLUS NEWS 35 Apr 21 New current-awareness alert (SDI) frequency in WPIDS/WPINDEX/WPIX NEWS 36 Apr 28 RDISCLOSURE now available on STN NEWS EXPRESS April 4 CURRENT WINDOWS VERSION IS V6.01a, CURRENT MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP), AND CURRENT DISCOVER FILE IS DATED 01 APRIL 2003 NEWS HOURS STN Operating Hours Plus Help Desk Availability NEWS INTER General Internet Information NEWS LOGIN Welcome Banner and News Items

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FILE 'HOME' ENTERED AT 08:11:38 ON 05 MAY 2003

=> fil reg COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FILL ESTIMATED COST

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STRUCTURE FILE UPDATES: 2 MAY 2003 HIGHEST RN 509953-09-7
DICTIONARY FILE UPDATES: 2 MAY 2003 HIGHEST RN 509953-09-7

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

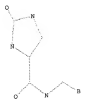
Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/Stnotes27.pdf

=> Uploading 10015328.str

L1 STRUCTURE UPLOADED

=> d L1 HAS NO ANSWERS L1 ST



Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 08:12:05 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -1 TO ITERATE

1 ITERATIONS 100.0% PROCESSED

1 ANSWERS

SEARCH TIME: 00.00.01

PROJECTED ANSWERS:

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE** 80 1 TO PROJECTED ITERATIONS: 1 TO 80

1 SEA SSS SAM L1

=> s 11 full

1.3

FULL SEARCH INITIATED 08:12:08 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 24 TO ITERATE

24 ITERATIONS 100.0% PROCESSED SEARCH TIME: 00.00.01

22 ANSWERS

22 SEA SSS FUL L1

=> fil caplus COST IN U.S. DOLLARS

TOTAL SINCE FILE SESSION ENTRY 148.36 148.15

FULL ESTIMATED COST

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FILE COVERS 1907 - 5 May 2003 VOL 138 ISS 19 FILE LAST UPDATED: 4 May 2003 (20030504/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 13

L4 1 L3

=> d ibib abs hitstr

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LA AMENIER 1 OF 1 I ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: INVENTOR(5): PATENT ASSIGNEE(5): SOUNCE: DOCUMENT TYPE: LANGUAGE: PAMILY ACC. NUM. COU	2002:466017 137:47198 Inddazoladan Hepatitis C v Han, Qi Bristol-Myers PCT Int. Appl CODEN: PIXXO Patent English	CAPLUS ines and their relat irrus NS3 protease : Squibb Pharma Com i., 173 pp.	.mhibitors
PATENT INFORMATION:		APPLICATION N	naTE
Wn 2002048152	A2 20020620	WO 2001-US479	16 20011212
W: AE, AG,	AL, AM, AT, AU, A	AZ, BA, BB, BG, BR,	BY, BZ, CA, CH, CN,
co, cx,	CU, CZ, DE, DX,	M, D2, EC, EE, ES,	FI, GB, GD, GE, GH,
GM, MR,	MU, ID, IL, IN,	IS, JP, KE, KG, KF,	KR, KZ, LC, LK, 1R, HZ, NO, HZ, PH, Pl.
LS, LT,	LU, LV, MA, ND,	4G, MX, HN, HW, MX,	TR, TT, TZ, UA, UG,
PT, RO.	RU, SD, SE, SO,	NZ, BY, KG, KE, HD,	DII 7.1 TM
U2, VN,	YU, ZA, ZN, AN,	20, 51, 60, 60, 110,	EM, EV, AT, BE, CH,
CI, DE,	DK, ED, F1, FA,	TA CHI GO GW MI	MR, NE, SN, TD, TG
85, 84,	CF, CG, CE, CA,	AU 2002-30764	20011212
AU 2002030764	AS LOUZDOLE		
PRIORITY APPLN. INFO	**	WO 2001-US47916	W 20011212
OTHER SOURCE(S):	MARPAT 137:4	7198	

- Title computs, were prepd. for use as serime protease inhibitors, emp.
 Hepatitis C virus NSD procease inhibitors (so data). Thus, the
 sundapolidinose I was obtained fore (8)-1-N-binarylowged tomyl-2-oxo-5inideolidinecarbowylic acid and the dioxaborolane fragment in 8 steps.
 47755-72-19 47735-73-29
- ANSWER 1 OF 1 CAPLUS COPPRIGHT 2003 ACS (Continued) 43755-43-6 CAPLUS COPPRIGHT 2003 ACS (Continued) 43755-43-6 CAPLUS CA

Absolute stereochemistry.

437153-44-7 (DADUS Carbon Land (15)-1-[(58)-5-[([(18)-1-(186,45,66,7a8)-hexablydro-channe and (15)-1-([(58)-5-[([(18)-1-(186,46,66,7a8)-hexablydro-channe (15)-6-(186)

Absolute stereochemistry.

437755-65-9 CAPLUS
4-Indazolidinecerboxamide, N-{(IR)-1-[(]38,48,65,78R)-hexahydro-18,5,5-transhyl-f-6-enshano-1,3,2-henzodioxaborol-2-y1]propyl-3-1-[(25)-2-enshyl-1-000-2-[([ghenylamno]cerbonyl]amno]butyl]-2-000-1-([-phenylamno]butyl)-4-(000-1-(-phenylamnoyl)-4-

Absolute stereochemistry

- AMERIA 1 of 1 CANAS COTTAINS 2003 ACS (CONLINSED)
 ALL EXT (Research) 500 Cyntains properties of 1 PME (Treparation) 5007
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Absolute stereochemistry.

Absolute stereochemistry.

- GTDELATE GTD

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS

437755-66-9 CAPLUS
4-Inidevolution search of the control of the co

43755-47-0 CAFLUS
4-Indisolidinecarbox manide, 3-([28]-2-[(9H-fluoren-1-ylcarboxy)]smino]-3-enchyl-1-nockyl)3-9-([1R]-1-[(2a],45,66,7ah)-hexalydic-3a,5,2-tramethyl-4,6-methano-1,3,2-bennodiceaborol-2-yl)propyl]-2-omo-1-(3-phenylpropyl)-((55)-(20))

Absolute stereochemistry.

L4 AMSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued)

FN 437755-48-1 CAPLUS
CM 4-Indiazolidianearboxanide, N-[(1R)-1-[(125,45,65,78R)-bexshydro-3s,5,5-trinstbyl-4,6-methano-1,3,2-benrodioxsborol-2-yl]sropyl-3-[(25)-2-[((4-methosyphenyl)acetyl]anisol-3-methyl-1-oxobutyl)-2-oxob-1-((3-phenylpropyl), ((5)-(CG)) (CA NIBOX MMS)

Absolute stereochemistry.

Appolute stereochemistry.

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued phenylpropyl)-, (48)- (901) (CA INDEX NAME)

N 43755-52-7 CAPLUS CM 4-Indiagolidamentobanide, 3-[(25)-2-[([1,1]-biphenyl]-4-ylaethyl]amino]-3-esthyl-1-osobutyl]-8-[([1,1]-1-[(3a5,45,65,7ah)-hexahydro-3a,5,5-trimethyl-4,6-esthano-1,3-2-benroi osobool-2-yl]propyl)-2-oxo-1-[1-phenyl-propyl]-, (45)-[02] [CA 18003 MMM]

Absolute stereochemistry.

38 43735-53-8 CAPAUS CX Carbance and, [1(3)-1-[1(3)-1-5][1(10)-1-1(3)-5,45,45,45,45] hexahydro-Ja,5,1-Elmandy L. bomming-1-2, based on backet and the comtraction of the community of the community of the community of the land of the community of the commun

Absolute stereochemistry.

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued)

39 437785-50-5 CM2UT CX Carbonac sed. [133-1-[1(55)-5-[[[(18)-1-[1285,45,65,788]-bexabyirotropylete and pleid, 6-sethano-1,3,2-bearodd seaborol-2ylpegogland on carboyl-12-ono-1-(3-phonylpropyl)-1 madselidacyl carboyl)-2-sethylpropyl-, 99:fluoren-9-ylsethyl ester [9CI] (CX LDEX MANS)

Absolute stereochemistry.

3N 437754-51-6 CAPJUS CN 4-Imidazolidinecarboxamide, N-[(1R)-1-[(385,45,65,7aR)-hexahydro-3a,5,5-true:hyl-4,6-enthano-1,3,2-henzodiuxaborol-2-yl]propyl)-3-[[28]-3-methyl-1-cox-2-([3-truit-thuo-comethyl])methyl]amino]butyl)-2-oxo-1-(3-truit-thuo-comethyl)methyl]amino]butyl)-2-oxo-1-(3-truit-thuo-comethyl)methyl]amino]butyl)-2-oxo-1-(3-truit-thuo-comethyl)methyl]amino]butyl)-3-([3-truit-thuo-comethyl)methyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-comethyllamino]butyl)-3-([3-truit-thuo-come

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued)

RW 437755-54-9 CAPLES
CN 3-Pyridisecarboxande6, 2-chloro-N=[(15)-1-[(15)-5-[[(110)-1[(136,46,67,32]-beashydro-3a,5,5-trinetbyl-4,6-sethano-1,3,2bensodiosaborol-2-yllproyyllanisolcarboxyll-2-oso-3-[3-phenylprosyyl-imidzabliosyl[carboxyll-2-entbylproyyl]-[051](CN INDEX MORE)

Absolute stereochemistry.

38 437755-55-0 CAPLUS ON 4-Inidacolidine-aboxanide, 3-[(25)-2-[(4-butylbantoy2) anino]-3-methyl-1-ocobutyl-1-fi-([35]-4,6-6,783]-berskhyl-7-5,5,5-frianthyl-4,6-nethano-1,3,2-bensodioxaborol-2-yi]propyl]-2-oco-1-(3-phanylpropyl)-, ([45]-(85)] (CA) 1802-3806

Absolute etereochemistry

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued)

437155-56-1 CMFLUS
Garbanic scid, [[15]-1-[[15]-5-[[[18]-1-[18,45,65,78h]-beaubydroGarbanic scid, [[15]-1-[15]-1-[18,45,65,78h]-beaubydroJillogovi lana, octobovi)-2-cos-3-[3-behydphogovi]-1saidacilisy||carbony||-2-sethy|propy||-, 2-sethy|propy| seter [9CI| [CA
INDOX DMS]]

Absolute stereochemistry.

437755-57-2 CAPLUS
4-[midzolidinecarbowanade, 3-[(25)-2-[[(benzylanino)carbonyi]amino]-3mathyl-1-oxebyuy]-N-[(1R)-1-[(3a5,45,65,7aR)-hexabydro-3a,5,5-t-timetbyl4,6-sethamo-1,3,2-benzeddioxeborol-2-yl]propyl]-2-oxe-1-(3-phenylpropyl)-,
(65)-[(21)-(CA INDEX NAME)

Absolute stereochemistry.

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued)

437755-60-7 CAPLUS
4-Inidazolidinecarboxamande, 3-[(25)-2-(benzoylamino)-3-nethyl-1-oxobutyl]-N-[(18)-1-(184,46,45,7a3)-beaxblydro-3a,5,5-trizentyl-4,6-methano-1,3,2-beaxod oxaborol-2-yl]propyl]-2-oxo-1-(3-phenylpropyl)-, (45)- (9CI) (CA HODEN MANN)

Absolute stereochemistry

437755-61-8 CAFLMS
1-leadsrolleinecarbonylo acad, 5-[[[[R]-1-[(3s.4s,65,65,7ak]-bagabydro-3a,5.*-translp1-4.6-estlamo-1,3,2-bascodowaborol-2-y1]-3-butenyljamnolocarbonyli-2-owos-3-[(227-3-phssyl-2-propenyl]-, phenylmethyl ester, (35) - (501 (CA INGEN INME)

Absolute stereochemistry. Double bond geometry as shown.

14 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued)

43755-58-3 CAPLUS
4-IndaScolid necarboxanide, N-[(IR)-1-[(Ja5,45,65,7aR)-hexabydro-3a,5,5-tranebly4-6,enebano-1,3,2-hearodioxaboro1-2-y1]propy1]-2-((Z5)-2-enby1-2-(L1)-apphthaliany) carboxy1 manno)-1-orobuty1]-2-oxo-1-(3-pheny)propy1]-, (45)- (471 (C4) TRUES MARE)

Absolute stereochemistry.

437755-59-4 CAPLUS
4-InidealidinearBowanide, 3-[(25)-2-(acetylamino)-3-methyl-1-oxobutyl]-N[(3)-1-(33, 45, 45, 45, 41)-harahydro-3a, 5, 5-ttmethyl-4, 6-methano-1, 3, 2-bencoinceaborol-2-yl]propyl]-2-oxo-1-(3-phenylpropyl)-, (45)-(56) (CA
ROOK ANNEL)

Absolute stereochemistry.

- L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS (Continued)
- RM 437785-62-9 CAPLIS CM 1-Indiagoldissecatopylic acid, S-{{{(IR)-1-{(IRS, 45, 65, 65, 788)-hexahydro-Ja, 5, 5-(raseby)-4-6-octbaso-{1,3,2-berzodioxaboro-{2-y1}-3-butsnyl]amino|actboxyl|-2-occ-3-{2-occ-2-(phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|ethyl|-phenylamino|

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